CALAN SR- verapamil hydrochloride tablet, film coated, extended release Pfizer Laboratories Div Pfizer Inc

CALAN® SR (Verapamil Hydrochloride) Extended-Release Tablets

DESCRIPTION

CALAN SR (verapamil hydrochloride) is a calcium ion influx inhibitor (slow-channel blocker or calcium ion antagonist). CALAN SR is available for oral administration as light green, capsule-shaped, scored, film-coated tablets containing 240 mg of verapamil hydrochloride (equivalent to 222.16 mg verapamil free base); as light pink, oval, scored, film-coated tablets containing 180 mg of verapamil hydrochloride (equivalent to 166.62 mg verapamil free base); and as light violet, oval, film-coated tablets containing 120 mg of verapamil hydrochloride (equivalent to 111.08 mg verapamil free base). The tablets are designed for extended-release of the drug in the gastrointestinal tract; extended-release characteristics are not altered when the tablet is divided in half.

The structural formula of verapamil HCl is:

$$\begin{array}{c} \text{CH}_3\text{O} & \text{CH}_3 \\ \text{CH}_3\text{O} & \text{CH}_3 \\ \text{C}_1\text{CH}_2\text{)}_3\text{NCH}_2\text{CH}_2 \\ \text{CH}_1\text{CH}_3\text{)}_2 \\ \text{C}_{27}\text{H}_{38}\text{N}_2\text{O}_4 \\ \end{array} \\ \text{HCl} & \text{M.W.=491.08} \\ \end{array}$$

Benzeneacetonitrile, α-[3-[[2-(3,4-dimethoxyphenyl)ethyl] methylamino]propyl]-3,4-dimethoxy-α-(1-methylethyl) hydrochloride

Verapamil HCl is an almost white, crystalline powder, practically free of odor, with a bitter taste. It is soluble in water, chloroform, and methanol. Verapamil HCl is not chemically related to other cardioactive drugs.

Inactive ingredients include alginate, carnauba wax, hypromellose, magnesium stearate, microcrystalline cellulose, polyethylene glycol, polyvinyl pyrrolidone, talc, titanium dioxide, and coloring agents: 240 mg—D&C Yellow No. 10 Lake and FD&C Blue No. 2 Lake; 120 and 180 mg—iron oxide.

CLINICAL PHARMACOLOGY

CALAN (verapamil HCl) is a calcium ion influx inhibitor (slow-channel blocker or calcium ion antagonist) that exerts its pharmacologic effects by modulating the influx of ionic calcium across the cell membrane of the arterial smooth muscle as well as in conductile and contractile myocardial cells.

Mechanism of action

Essential hypertension

Verapamil exerts antihypertensive effects by decreasing systemic vascular resistance, usually without orthostatic decreases in blood pressure or reflex tachycardia; bradycardia (rate less than 50 beats/min) is uncommon (1.4%). During isometric or dynamic exercise, CALAN does not alter systolic cardiac

function in patients with normal ventricular function.

CALAN does not alter total serum calcium levels. However, one report suggested that calcium levels above the normal range may alter the therapeutic effect of CALAN.

Other pharmacologic actions of CALAN include the following

CALAN dilates the main coronary arteries and coronary arterioles, both in normal and ischemic regions, and is a potent inhibitor of coronary artery spasm, whether spontaneous or ergonovine-induced. This property increases myocardial oxygen delivery in patients with coronary artery spasm and is responsible for the effectiveness of CALAN in vasospastic (Prinzmetal's or variant) as well as unstable angina at rest. Whether this effect plays any role in classical effort angina is not clear, but studies of exercise tolerance have not shown an increase in the maximum exercise rate—pressure product, a widely accepted measure of oxygen utilization. This suggests that, in general, relief of spasm or dilation of coronary arteries is not an important factor in classical angina.

CALAN regularly reduces the total systemic resistance (afterload) against which the heart works both at rest and at a given level of exercise by dilating peripheral arterioles.

Electrical activity through the AV node depends, to a significant degree, upon calcium influx through the slow channel. By decreasing the influx of calcium, CALAN prolongs the effective refractory period within the AV node and slows AV conduction in a rate-related manner.

Normal sinus rhythm is usually not affected, but in patients with sick sinus syndrome, CALAN may interfere with sinus-node impulse generation and may induce sinus arrest or sinoatrial block. Atrioventricular block can occur in patients without preexisting conduction defects (see **WARNINGS**).

CALAN does not alter the normal atrial action potential or intraventricular conduction time, but depresses amplitude, velocity of depolarization, and conduction in depressed atrial fibers. CALAN may shorten the antegrade effective refractory period of the accessory bypass tract. Acceleration of ventricular rate and/or ventricular fibrillation has been reported in patients with atrial flutter or atrial fibrillation and a coexisting accessory AV pathway following administration of verapamil (see **WARNINGS**).

CALAN has a local anesthetic action that is 1.6 times that of procaine on an equimolar basis. It is not known whether this action is important at the doses used in man.

Pharmacokinetics and metabolism

With the immediate-release formulation, more than 90% of the orally administered dose of CALAN is absorbed. Because of rapid biotransformation of verapamil during its first pass through the portal circulation, bioavailability ranges from 20% to 35%. Peak plasma concentrations are reached between 1 and 2 hours after oral administration. Chronic oral administration of 120 mg of verapamil HCl every 6 hours resulted in plasma levels of verapamil ranging from 125 to 400 ng/mL, with higher values reported occasionally. A nonlinear correlation between the verapamil dose administered and verapamil plasma level does exist. In early dose titration with verapamil, a relationship exists between verapamil plasma concentration and prolongation of the PR interval. However, during chronic administration this relationship may disappear. The mean elimination half-life in single-dose studies ranged from 2.8 to 7.4 hours. In these same studies, after repetitive dosing, the half-life increased to a range from 4.5 to 12.0 hours (after less than 10 consecutive doses given 6 hours apart). Half-life of verapamil may increase during titration. No relationship has been established between the plasma concentration of verapamil and a reduction in blood pressure.

Aging may affect the pharmacokinetics of verapamil. Elimination half-life may be prolonged in the elderly. In multiple-dose studies under fasting conditions, the bioavailability, measured by AUC, of CALAN SR was similar to CALAN (immediate release); rates of absorption were of course different.

In a randomized, single-dose, crossover study using healthy volunteers, administration of 240 mg CALAN SR with food produced peak plasma verapamil concentrations of 79 ng/mL; time to peak plasma

verapamil concentration of 7.71 hours; and AUC (0–24 hr) of 841 ng·hr/mL. When CALAN SR was administered to fasting subjects, peak plasma verapamil concentration was 164 ng/mL; time to peak plasma verapamil concentration was 5.21 hours; and AUC (0–24 hr) was 1,478 ng·hr/mL. Similar results were demonstrated for plasma norverapamil. Food thus produces decreased bioavailability (AUC) but a narrower peak-to-trough ratio. Good correlation of dose and response is not available, but controlled studies of CALAN SR have shown effectiveness of doses similar to the effective doses of CALAN (immediate release).

In healthy men, orally administered CALAN undergoes extensive metabolism in the liver. Twelve metabolites have been identified in plasma; all except norverapamil are present in trace amounts only. Norverapamil can reach steady-state plasma concentrations approximately equal to those of verapamil itself. The cardiovascular activity of norverapamil appears to be approximately 20% that of verapamil. Approximately 70% of an administered dose is excreted as metabolites in the urine and 16% or more in the feces within 5 days. About 3% to 4% is excreted in the urine as unchanged drug. Approximately 90% is bound to plasma proteins. In patients with hepatic insufficiency, metabolism of immediate-release verapamil is delayed and elimination half-life prolonged up to 14 to 16 hours (see **PRECAUTIONS**); the volume of distribution is increased and plasma clearance reduced to about 30% of normal. Verapamil clearance values suggest that patients with liver dysfunction may attain therapeutic verapamil plasma concentrations with one third of the oral daily dose required for patients with normal liver function.

After four weeks of oral dosing (120 mg q.i.d.), verapamil and norverapamil levels were noted in the cerebrospinal fluid with estimated partition coefficient of 0.06 for verapamil and 0.04 for norverapamil.

In ten healthy males, administration of oral verapamil (80 mg every 8 hours for 6 days) and a single oral dose of ethanol (0.8 g/kg) resulted in a 17% increase in mean peak ethanol concentrations (106.45 \pm 21.40 to 124.23 \pm 24.74 mg·hr/dL) compared to placebo. The area under the blood ethanol concentration versus time curve (AUC over 12 hours) increased by 30% (365.67 \pm 93.52 to 475.07 \pm 97.24 mg·hr/dL). Verapamil AUCs were positively correlated (r=0.71) to increased ethanol blood AUC values (see **PRECAUTIONS, Drug interactions**).

Hemodynamics and myocardial metabolism

CALAN reduces afterload and myocardial contractility. Improved left ventricular diastolic function in patients with Idiopathic Hypertrophic Subaortic Stenosis (IHSS) and those with coronary heart disease has also been observed with CALAN. In most patients, including those with organic cardiac disease, the negative inotropic action of CALAN is countered by reduction of afterload, and cardiac index is usually not reduced. However, in patients with severe left ventricular dysfunction (eg, pulmonary wedge pressure above 20 mm Hg or ejection fraction less than 30%), or in patients taking beta-adrenergic blocking agents or other cardiodepressant drugs, deterioration of ventricular function may occur (see **PRECAUTIONS, Drug interactions**).

Pulmonary function

CALAN does not induce bronchoconstriction and, hence, does not impair ventilatory function.

INDICATIONS AND USAGE

CALAN SR is indicated for the treatment of hypertension, to lower blood pressure. Lowering blood pressure reduces the risk of fatal and nonfatal cardiovascular events, primarily strokes and myocardial infarctions. These benefits have been seen in controlled trials of antihypertensive drugs from a wide variety of pharmacologic classes including this drug.

Control of high blood pressure should be part of comprehensive cardiovascular risk management, including, as appropriate, lipid control, diabetes management, antithrombotic therapy, smoking cessation,

exercise, and limited sodium intake. Many patients will require more than one drug to achieve blood pressure goals. For specific advice on goals and management, see published guidelines, such as those of the National High Blood Pressure Education Program's Joint National Committee on Prevention, Detection, Evaluation, and Treatment of High Blood Pressure (JNC).

Numerous antihypertensive drugs, from a variety of pharmacologic classes and with different mechanisms of action, have been shown in randomized controlled trials to reduce cardiovascular morbidity and mortality, and it can be concluded that it is blood pressure reduction, and not some other pharmacologic property of the drugs, that is largely responsible for those benefits. The largest and most consistent cardiovascular outcome benefit has been a reduction in the risk of stroke, but reductions in myocardial infarction and cardiovascular mortality also have been seen regularly.

Elevated systolic or diastolic pressure causes increased cardiovascular risk, and the absolute risk increase per mmHg is greater at higher blood pressures, so that even modest reductions of severe hypertension can provide substantial benefit. Relative risk reduction from blood pressure reduction is similar across populations with varying absolute risk, so the absolute benefit is greater in patients who are at higher risk independent of their hypertension (for example, patients with diabetes or hyperlipidemia), and such patients would be expected to benefit from more aggressive treatment to a lower blood pressure goal.

Some antihypertensive drugs have smaller blood pressure effects (as monotherapy) in black patients, and many antihypertensive drugs have additional approved indications and effects (e.g., on angina, heart failure, or diabetic kidney disease). These considerations may guide selection of therapy.

CONTRAINDICATIONS

Verapamil HCl extended-release tablets are contraindicated in:

- 1. Severe left ventricular dysfunction (see **WARNINGS**)
- 2. Hypotension (systolic pressure less than 90 mm Hg) or cardiogenic shock
- 3. Sick sinus syndrome (except in patients with a functioning artificial ventricular pacemaker)
- 4. Second- or third-degree AV block (except in patients with a functioning artificial ventricular pacemaker)
- 5. Patients with atrial flutter or atrial fibrillation and an accessory bypass tract (eg, Wolff-Parkinson-White, Lown-Ganong-Levine syndromes) (see **WARNINGS**)
- 6. Patients with known hypersensitivity to verapamil hydrochloride

WARNINGS

Heart failure

Verapamil has a negative inotropic effect, which in most patients is compensated by its afterload reduction (decreased systemic vascular resistance) properties without a net impairment of ventricular performance. In clinical experience with 4,954 patients, 87 (1.8%) developed congestive heart failure or pulmonary edema. Verapamil should be avoided in patients with severe left ventricular dysfunction (eg, ejection fraction less than 30%) or moderate to severe symptoms of cardiac failure and in patients with any degree of ventricular dysfunction if they are receiving a beta-adrenergic blocker (see **PRECAUTIONS**, **Drug interactions**). Patients with milder ventricular dysfunction should, if possible, be controlled with optimum doses of digitalis and/or diuretics before verapamil treatment. (**Note interactions with digoxin under PRECAUTIONS**)

Hypotension

Occasionally, the pharmacologic action of verapamil may produce a decrease in blood pressure below normal levels, which may result in dizziness or symptomatic hypotension. The incidence of hypotension observed in 4,954 patients enrolled in clinical trials was 2.5%. In hypertensive patients, decreases in

blood pressure below normal are unusual. Tilt-table testing (60 degrees) was not able to induce orthostatic hypotension.

Elevated liver enzymes

Elevations of transaminases with and without concomitant elevations in alkaline phosphatase and bilirubin have been reported. Such elevations have sometimes been transient and may disappear even in the face of continued verapamil treatment. Several cases of hepatocellular injury related to verapamil have been proven by rechallenge; half of these had clinical symptoms (malaise, fever, and/or right upper quadrant pain) in addition to elevation of SGOT, SGPT, and alkaline phosphatase. Periodic monitoring of liver function in patients receiving verapamil is therefore prudent.

Accessory bypass tract (Wolff-Parkinson-White or Lown-Ganong-Levine)

Some patients with paroxysmal and/or chronic atrial fibrillation or atrial flutter and a coexisting accessory AV pathway have developed increased antegrade conduction across the accessory pathway bypassing the AV node, producing a very rapid ventricular response or ventricular fibrillation after receiving intravenous verapamil (or digitalis). Although a risk of this occurring with oral verapamil has not been established, such patients receiving oral verapamil may be at risk and its use in these patients is contraindicated (see **CONTRAINDICATIONS**). Treatment is usually DC-cardioversion. Cardioversion has been used safely and effectively after oral CALAN.

Atrioventricular block

The effect of verapamil on AV conduction and the SA node may cause asymptomatic first-degree AV block and transient bradycardia, sometimes accompanied by nodal escape rhythms. PR-interval prolongation is correlated with verapamil plasma concentrations, especially during the early titration phase of therapy. Higher degrees of AV block, however, were infrequently (0.8%) observed. Marked first-degree block or progressive development to second- or third-degree AV block, requires a reduction in dosage or, in rare instances, discontinuation of verapamil HCl and institution of appropriate therapy, depending upon the clinical situation.

Patients with hypertrophic cardiomyopathy (IHSS)

In 120 patients with hypertrophic cardiomyopathy (most of them refractory or intolerant to propranolol) who received therapy with verapamil at doses up to 720 mg/day, a variety of serious adverse effects were seen. Three patients died in pulmonary edema; all had severe left ventricular outflow obstruction and a past history of left ventricular dysfunction. Eight other patients had pulmonary edema and/or severe hypotension; abnormally high (greater than 20 mm Hg) pulmonary wedge pressure and a marked left ventricular outflow obstruction were present in most of these patients. Concomitant administration of quinidine (see **PRECAUTIONS, Drug interactions**) preceded the severe hypotension in 3 of the 8 patients (2 of whom developed pulmonary edema). Sinus bradycardia occurred in 11% of the patients, second-degree AV block in 4%, and sinus arrest in 2%. It must be appreciated that this group of patients had a serious disease with a high mortality rate. Most adverse effects responded well to dose reduction, and only rarely did verapamil use have to be discontinued.

PRECAUTIONS

General

Use in patients with impaired hepatic function

Since verapamil is highly metabolized by the liver, it should be administered cautiously to patients with impaired hepatic function. Severe liver dysfunction prolongs the elimination half-life of immediate-release verapamil to about 14 to 16 hours; hence, approximately 30% of the dose given to patients with normal liver function should be administered to these patients. Careful monitoring for abnormal

prolongation of the PR interval or other signs of excessive pharmacologic effects (see **OVERDOSAGE**) should be carried out.

Use in patients with attenuated (decreased) neuromuscular transmission

It has been reported that verapamil decreases neuromuscular transmission in patients with Duchenne's muscular dystrophy, and that verapamil prolongs recovery from the neuromuscular blocking agent vecuronium. It may be necessary to decrease the dosage of verapamil when it is administered to patients with attenuated neuromuscular transmission.

Use in patients with impaired renal function

About 70% of an administered dose of verapamil is excreted as metabolites in the urine. Verapamil is not removed by hemodialysis. Until further data are available, verapamil should be administered cautiously to patients with impaired renal function. These patients should be carefully monitored for abnormal prolongation of the PR interval or other signs of overdosage (see **OVERDOSAGE**).

Drug interactions

Cytochrome inducers/inhibitors

In vitro metabolic studies indicate that verapamil is metabolized by cytochrome P450 CYP3A4, CYP1A2, CYP2C8, CYP2C9, and CYP2C18. Clinically significant interactions have been reported with inhibitors of CYP3A4 (e.g., erythromycin, ritonavir) causing elevation of plasma levels of verapamil while inducers of CYP3A4 (e.g., rifampin) have caused a lowering of plasma levels of verapamil.

HMG-CoA reductase inhibitors

The use of HMG-CoA reductase inhibitors that are CYP3A4 substrates in combination with verapamil has been associated with reports of myopathy/rhabdomyolysis.

Co-administration of multiple doses of 10 mg of verapamil with 80 mg simvastatin resulted in exposure to simvastatin 2.5-fold that following simvastatin alone. Limit the dose of simvastatin in patients on verapamil to 10 mg daily. Limit the daily dose of lovastatin to 40 mg. Lower starting and maintenance doses of other CYP3A4 substrates (e.g., atorvastatin) may be required as verapamil may increase the plasma concentration of these drugs.

Ivabradine

Concurrent use of verapamil increases exposure to ivabradine and may exacerbate bradycardia and conduction disturbances. Avoid co-administration of verapamil and ivabradine.

Beta-blockers

Concomitant therapy with beta-adrenergic blockers and verapamil may result in additive negative effects on heart rate, atrioventricular conduction and/or cardiac contractility. The combination of extended-release verapamil and beta-adrenergic blocking agents has not been studied. However, there have been reports of excessive bradycardia and AV block, including complete heart block, when the combination has been used for the treatment of hypertension. For hypertensive patients, the risks of combined therapy may outweigh the potential benefits. The combination should be used only with caution and close monitoring.

Asymptomatic bradycardia (36 beats/min) with a wandering atrial pacemaker has been observed in a patient receiving concomitant timolol (a beta-adrenergic blocker) eyedrops and oral verapamil.

A decrease in metoprolol and propranolol clearance has been observed when either drug is administered concomitantly with verapamil. A variable effect has been seen when verapamil and atenolol were given together.

Digitalis

Clinical use of verapamil in digitalized patients has shown the combination to be well tolerated if digoxin doses are properly adjusted. However, chronic verapamil treatment can increase serum digoxin levels by 50% to 75% during the first week of therapy, and this can result in digitalis toxicity. In patients with hepatic cirrhosis the influence of verapamil on digoxin kinetics is magnified. Verapamil may reduce total body clearance and extrarenal clearance of digitoxin by 27% and 29%, respectively. Maintenance and digitalization doses should be reduced when verapamil is administered, and the patient should be carefully monitored to avoid over- or under-digitalization. Whenever over-digitalization is suspected, the daily dose of digitalis should be reduced or temporarily discontinued. On discontinuation of CALAN use, the patient should be reassessed to avoid under-digitalization.

Antihypertensive agents

Verapamil administered concomitantly with oral antihypertensive agents (eg, vasodilators, angiotensin-converting enzyme inhibitors, diuretics, beta-blockers) will usually have an additive effect on lowering blood pressure. Patients receiving these combinations should be appropriately monitored. Concomitant use of agents that attenuate alpha-adrenergic function with verapamil may result in a reduction in blood pressure that is excessive in some patients. Such an effect was observed in one study following the concomitant administration of verapamil and prazosin.

Antiarrhythmic agents

Disopyramide

Until data on possible interactions between verapamil and disopyramide phosphate are obtained, disopyramide should not be administered within 48 hours before or 24 hours after verapamil administration.

Flecainide

A study in healthy volunteers showed that the concomitant administration of flecainide and verapamil may have additive effects on myocardial contractility, AV conduction, and repolarization. Concomitant therapy with flecainide and verapamil may result in additive negative inotropic effect and prolongation of atrioventricular conduction.

Quinidine

In a small number of patients with hypertrophic cardiomyopathy (IHSS), concomitant use of verapamil and quinidine resulted in significant hypotension. Until further data are obtained, combined therapy of verapamil and quinidine in patients with hypertrophic cardiomyopathy should probably be avoided.

The electrophysiologic effects of quinidine and verapamil on AV conduction were studied in 8 patients. Verapamil significantly counteracted the effects of quinidine on AV conduction. There has been a report of increased quinidine levels during verapamil therapy.

Other agents

Alcohol

Verapamil has been found to inhibit ethanol elimination significantly, resulting in elevated blood ethanol concentrations that may prolong the intoxicating effects of alcohol (see **CLINICAL**

PHARMACOLOGY, Pharmacokinetics and metabolism).

Nitrates

Verapamil has been given concomitantly with short- and long-acting nitrates without any undesirable drug interactions. The pharmacologic profile of both drugs and the clinical experience suggest beneficial interactions.

Cimetidine

The interaction between cimetidine and chronically administered verapamil has not been studied. Variable results on clearance have been obtained in acute studies of healthy volunteers; clearance of verapamil was either reduced or unchanged.

Lithium

Increased sensitivity to the effects of lithium (neurotoxicity) has been reported during concomitant verapamil-lithium therapy; lithium levels have been observed sometimes to increase, sometimes to decrease, and sometimes to be unchanged. Patients receiving both drugs must be monitored carefully.

Carbamazepine

Verapamil therapy may increase carbamazepine concentrations during combined therapy. This may produce carbamazepine side effects such as diplopia, headache, ataxia, or dizziness.

Rifampin

Therapy with rifampin may markedly reduce oral verapamil bioavailability.

Phenobarbital

Phenobarbital therapy may increase verapamil clearance.

Cyclosporin

Verapamil therapy may increase serum levels of cyclosporin.

Theophylline

Verapamil may inhibit the clearance and increase the plasma levels of theophylline.

Inhalation anesthetics

Animal experiments have shown that inhalation anesthetics depress cardiovascular activity by decreasing the inward movement of calcium ions. When used concomitantly, inhalation anesthetics and calcium antagonists, such as verapamil, should each be titrated carefully to avoid excessive cardiovascular depression.

Neuromuscular blocking agents

Clinical data and animal studies suggest that verapamil may potentiate the activity of neuromuscular blocking agents (curare-like and depolarizing). It may be necessary to decrease the dose of verapamil and/or the dose of the neuromuscular blocking agent when the drugs are used concomitantly.

Telithromycin

Hypotension and bradyarrhythmias have been observed in patients receiving concurrent telithromycin, an antibiotic in the ketolide class.

Clonidine

Sinus bradycardia resulting in hospitalization and pacemaker insertion has been reported in association with the use of clonidine concurrently with verapamil. Monitor heart rate in patients receiving concomitant verapamil and clonidine.

Mammalian target of rapamycin (mTOR) inhibitors

In a study of 25 healthy volunteers with co-administration of verapamil with sirolimus, whole blood

sirolimus C_{max} and AUC were increased 130% and 120%, respectively. Plasma S-(-) verapamil C_{max} and AUC were both increased 50%. Co-administration of verapamil with everolimus in 16 healthy volunteers increased the C_{max} and AUC of everolimus by 130% and 250%, respectively. With concomitant use of mTOR inhibitors (e.g., sirolimus, temsirolimus, and everolimus) and verapamil, consider appropriate dose reductions of both medications.

Carcinogenesis, mutagenesis, impairment of fertility

An 18-month toxicity study in rats, at a low multiple (6-fold) of the maximum recommended human dose, and not the maximum tolerated dose, did not suggest a tumorigenic potential. There was no evidence of a carcinogenic potential of verapamil administered in the diet of rats for two years at doses of 10, 35, and 120 mg/kg/day or approximately 1, 3.5, and 12 times, respectively, the maximum recommended human daily dose (480 mg/day or 9.6 mg/kg/day).

Verapamil was not mutagenic in the Ames test in 5 test strains at 3 mg per plate with or without metabolic activation.

Studies in female rats at daily dietary doses up to 5.5 times (55 mg/kg/day) the maximum recommended human dose did not show impaired fertility. Effects on male fertility have not been determined.

Pregnancy

Reproduction studies have been performed in rabbits and rats at oral doses up to 1.5 (15 mg/kg/day) and 6 (60 mg/kg/day) times the human oral daily dose, respectively, and have revealed no evidence of teratogenicity. In the rat, however, this multiple of the human dose was embryocidal and retarded fetal growth and development, probably because of adverse maternal effects reflected in reduced weight gains of the dams. This oral dose has also been shown to cause hypotension in rats. There are no adequate and well-controlled studies in pregnant women. Because animal reproduction studies are not always predictive of human response, this drug should be used during pregnancy only if clearly needed. Verapamil crosses the placental barrier and can be detected in umbilical vein blood at delivery.

Labor and delivery

It is not known whether the use of verapamil during labor or delivery has immediate or delayed adverse effects on the fetus, or whether it prolongs the duration of labor or increases the need for forceps delivery or other obstetric intervention. Such adverse experiences have not been reported in the literature, despite a long history of use of verapamil in Europe in the treatment of cardiac side effects of beta-adrenergic agonist agents used to treat premature labor.

Nursing mothers

Verapamil is excreted in human milk. Because of the potential for adverse reactions in nursing infants from verapamil, nursing should be discontinued while verapamil is administered.

Pediatric use

Safety and efficacy of CALAN SR in pediatric patients below the age of 18 years have not been established.

Animal pharmacology and/or animal toxicology

In chronic animal toxicology studies, verapamil caused lenticular and/or suture line changes at 30 mg/kg/day or greater, and frank cataracts at 62.5 mg/kg/day or greater in the beagle dog but not in the rat. Development of cataracts due to verapamil has not been reported in man.

ADVERSE REACTIONS

Serious adverse reactions are uncommon when verapamil therapy is initiated with upward dose titration

within the recommended single and total daily dose. See **WARNINGS** for discussion of heart failure, hypotension, elevated liver enzymes, AV block, and rapid ventricular response. Reversible (upon discontinuation of verapamil) non-obstructive, paralytic ileus has been infrequently reported in association with the use of verapamil. The following reactions to orally administered verapamil occurred at rates greater than 1.0% or occurred at lower rates but appeared clearly drug-related in clinical trials in 4,954 patients:

Constipation	7.3%	Dyspnea	1.4%
Dizziness	3.3%	Bradycardia	
Nausea	2.7%	(HR <50/min)	1.4%
Hypotension	2.5%	AV block	
Headache	2.2%	(total 1°, 2°, 3°)	1.2%
Edema	1.9%	(2° and 3°)	0.8%
CHF, Pulmonary edema	1.8%	Rash	1.2%
Fatigue	1.7%	Flushing	0.6%

Elevated liver enzymes (see **WARNINGS**)

In clinical trials related to the control of ventricular response in digitalized patients who had atrial fibrillation or flutter, ventricular rates below 50/min at rest occurred in 15% of patients and asymptomatic hypotension occurred in 5% of patients.

The following reactions, reported in 1% or less of patients, occurred under conditions (open trials, marketing experience) where a causal relationship is uncertain; they are listed to alert the physician to a possible relationship:

Cardiovascular: angina pectoris, atrioventricular dissociation, chest pain, claudication, myocardial infarction, palpitations, purpura (vasculitis), syncope.

Digestive system: diarrhea, dry mouth, gastrointestinal distress, gingival hyperplasia.

Hemic and lymphatic: ecchymosis or bruising.

Nervous system: cerebrovascular accident, confusion, equilibrium disorders, insomnia, muscle cramps, paresthesia, psychotic symptoms, shakiness, somnolence.

Skin: arthralgia and rash, exanthema, hair loss, hyperkeratosis, macules, sweating, urticaria, Stevens-Johnson syndrome, erythema multiforme.

Special senses: blurred vision, tinnitus.

Urogenital: gynecomastia, galactorrhea/hyperprolactinemia, increased urination, spotty menstruation, impotence.

Treatment of acute cardiovas cular adverse reactions

The frequency of cardiovascular adverse reactions that require therapy is rare; hence, experience with their treatment is limited. Whenever severe hypotension or complete AV block occurs following oral administration of verapamil, the appropriate emergency measures should be applied immediately; eg, intravenously administered norepinephrine bitartrate, atropine sulfate, isoproterenol HCl (all in the usual doses), or calcium gluconate (10% solution). In patients with hypertrophic cardiomyopathy (IHSS), alpha-adrenergic agents (phenylephrine HCl, metaraminol bitartrate, or methoxamine HCl) should be used to maintain blood pressure, and isoproterenol and norepinephrine should be avoided. If further support is necessary, dopamine HCl or dobutamine HCl may be administered. Actual treatment and dosage should depend on the severity of the clinical situation and the judgment and experience of the treating physician.

OVERDOSAGE

Overdose with verapamil may lead to pronounced hypotension, bradycardia, and conduction system abnormalities (eg, junctional rhythm with AV dissociation and high degree AV block, including asystole). Other symptoms secondary to hypoperfusion (eg, metabolic acidosis, hyperglycemia, hyperkalemia, renal dysfunction, and convulsions) may be evident.

Treat all verapamil overdoses as serious and maintain observation for at least 48 hours (especially CALAN SR), preferably under continuous hospital care. Delayed pharmacodynamic consequences may occur with the extended-release formulation. Verapamil is known to decrease gastrointestinal transit time.

In overdose, extended-release tablets of CALAN SR have occasionally been reported to form concretions within the stomach or intestines. These concretions have not been visible on plain radiographs of the abdomen, and no medical means of gastrointestinal emptying is of proven efficacy in removing them. Endoscopy might reasonably be considered in cases of massive overdose when symptoms are unusually prolonged.

Treatment of overdosage should be supportive. Beta-adrenergic stimulation or parenteral administration of calcium solutions may increase calcium ion flux across the slow channel and have been used effectively in treatment of deliberate overdosage with verapamil. Continued treatment with large doses of calcium may produce a response. In a few reported cases, overdose with calcium channel blockers that was initially refractory to atropine became more responsive to this treatment when the patients received large doses (close to 1 g/hr for more than 24 hr) of calcium chloride. Verapamil cannot be removed by hemodialysis. Clinically significant hypotensive reactions or high degree AV block should be treated with vasopressor agents or cardiac pacing, respectively. Asystole should be handled by the usual measures including cardiopulmonary resuscitation.

DOSAGE AND ADMINISTRATION

Essential hypertension

The dose of CALAN SR should be individualized by titration and the drug should be administered with food. Initiate therapy with 180 mg of extended-release verapamil HCl, CALAN SR, given in the morning. Lower initial doses of 120 mg a day may be warranted in patients who may have an increased response to verapamil (eg, the elderly or small people). Upward titration should be based on therapeutic efficacy and safety evaluated weekly and approximately 24 hours after the previous dose. The antihypertensive effects of CALAN SR are evident within the first week of therapy.

If adequate response is not obtained with 180 mg of CALAN SR, the dose may be titrated upward in the following manner:

- a) 240 mg each morning,
- b) 180 mg each morning plus
 - 180 mg each evening; or
 - 240 mg each morning plus
 - 120 mg each evening,
- c) 240 mg every 12 hours.

When switching from immediate-release CALAN to CALAN SR, the total daily dose in milligrams may remain the same.

HOW SUPPLIED

CALAN SR 120 mg extended-release tablets are light violet, oval, film coated, with CALAN debossed

on one side and SR 120 on the other, supplied as:

NDC Number	Size
0025-1901-31	bottle of
0023-1301-31	100
0025-2712-31	bottle of
0023-2/12-31	100

CALAN SR 180 mg extended-release tablets are light pink, oval, scored, film coated, with CALAN debossed on one side and SR 180 on the other, supplied as:

NDC Number	Size
0025-1911-31	bottle of
	100 bottle of
0025-2818-31	100

CALAN SR 240 mg extended-release tablets are light green, capsule shaped, scored, film coated, with CALAN debossed on one side and SR 240 on the other, supplied as:

NDC Number	Size
0025-1891-31	bottle of
0023-1031-31	100
0025-2924-31	bottle of
0025-2524-51	100

Store at 15° to 25° C (59° to 77° F) and protect from light and moisture. Dispense in tight, light-resistant containers.



LAB-0268-13.0 Revised March 2020

PRINCIPAL DISPLAY PANEL - 120 mg Caplet Bottle Label

NDC 0025-1901-31

Pfizer

Calan® SR

(verapamil

hydrochloride) caplets

SUSTAINED-RELEASE

120 mg

100 Caplets

Rx only



PRINCIPAL DISPLAY PANEL - 180 mg Caplet Bottle Label

NDC 0025-1911-31

Pfizer

Calan® SR

(verapamil

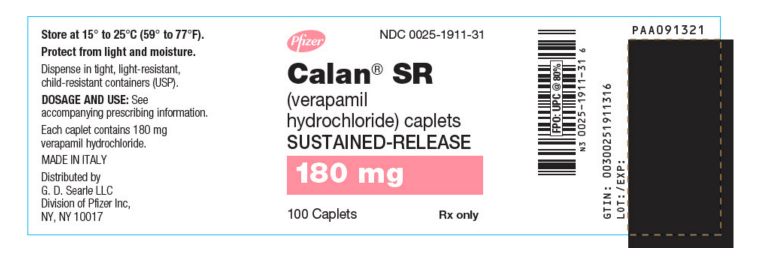
hydrochloride) caplets

SUSTAINED-RELEASE

180 mg

100 Caplets

Rx only



PRINCIPAL DISPLAY PANEL - 240 mg Caplet Bottle Label

NDC 0025-1891-31

Pfizer

Calan[®] SR

(verapamil

hydrochloride) caplets

SUSTAINED-RELEASE

240 mg

100 Caplets

Rx only



PRINCIPAL DISPLAY PANEL - 120 mg Tablet Bottle Label

NDC 0025-2712-31

Pfizer

Calan[®] SR (verapamil hydrochloride) extended-release tablets

120 mg*

100 Tablets

Rx only



PRINCIPAL DISPLAY PANEL - 180 mg Tablet Bottle Label

NDC 0025-2818-31 Pfizer Calan® SR

(verapamil hydrochloride) extended-release tablets

100 Tablets Rx only

Store at 15° to 25°C (59° to 77°F). Protect from light and moisture.

Dispense in tight, light-resistant, child-resistant containers (USP).

DOSAGE AND USE: See accompanying prescribing information.

*Each tablet contains verapamil hydrochloride 180 mg (equivalent to 166.62 mg verapamil).

MADE IN ITALY

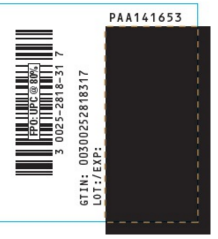
Distributed by G. D. Searle LLC Division of Pfizer Inc, NY, NY 10017 NDC 0025-2818-31

Calan® SR

(verapamil hydrochloride) extended-release tablets

180 mg*

100 Tablets Rx only



PRINCIPAL DISPLAY PANEL - 240 mg Tablet Bottle Label

NDC 0025-2924-31

Pfizer

Calan[®] SR (verapamil hydrochloride) extended-release tablets

240 mg*

100 Tablets

Rx only

Store at 15° to 25°C (59° to 77°F). Protect from light and moisture.

Dispense in tight, light-resistant, child-resistant containers (USP).

DOSAGE AND USE: See accompanying prescribing information.

*Each tablet contains verapamil hydrochloride 240 mg (equivalent to 222.16 mg verapamil).

MADE IN ITALY

Distributed by G. D. Searle LLC Division of Pfizer Inc, NY, NY 10017 Pfizer

NDC 0025-2924-31

Calan® SR

(verapamil hydrochloride) extended-release tablets

240 mg*

100 Tablets

Rx only



CALAN SR

verapamil hydrochloride tablet, film coated, extended release

Product Information

Product Type

HUMAN PRESCRIPTION DRUG

Item Code (Source)

NDC:0025-1901

ORAL

Active Ingredient/Active Moiety			
Ingredient Name	Basis of Strength	Strength	
VERAPAMIL HYDRO CHLO RIDE (UNII: V38880EY5R) (VERAPAMIL - UNII:CJ0 O 37 K U 29)	VERAPAMIL HYDROCHLORIDE	120 mg	

Inactive Ingredients		
Ingredient Name	Strength	
CARNAUBA WAX (UNII: R12CBM0 EIZ)		
HYPROMELLOSE, UNSPECIFIED (UNII: 3NXW29V3WO)		
MAGNESIUM STEARATE (UNII: 70097M6I30)		
MICRO CRYSTALLINE CELLULO SE (UNII: OP1R32D61U)		
POLYETHYLENE GLYCOL, UNSPECIFIED (UNII: 3WJQ0SDW1A)		
TALC (UNII: 7SEV7J4R1U)		
TITANIUM DIO XIDE (UNII: 15FIX9 V2JP)		

Product Characteristics			
Color	PURPLE (light violet)	Score	no score
Shape	OVAL	Size	11mm
Flavor		Imprint Code	CALAN;SR;120
Contains			

l	Packaging				
l	#	Item Code	Package Description	Marketing Start Date	Marketing End Date
	1	NDC:0025-1901-31	100 in 1 BOTTLE; Type 0: Not a Combination Product	03/06/1991	

Marketing Information			
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
NDA	NDA0 19 152	03/06/1991	

CALAN SR

Product Information				
Product Type	HUMAN PRESCRIPTION DRUG	Item Code (Source)	NDC:0025-1911	
Route of Administration	ORAL			

Active Ingredient/Active Moiety		
Ingredient Name	Basis of Strength	Strength

U U U U U U U U U U U U U U U U U U U		
VERAPAMIL HYDRO CHLO RIDE (UNII: V3888 O E Y5R) (VERAPAMIL -	VERAPAMIL	180 mg
UNII:CJ0O37KU29)	HYDROCHLORIDE	100 mg

Inactive Ingredients		
Ingredient Name	Strength	
CARNAUBA WAX (UNII: R12CBM0EIZ)		
HYPROMELLOSE, UNSPECIFIED (UNII: 3NXW29 V3WO)		
MAGNESIUM STEARATE (UNII: 70097M6I30)		
MICRO CRYSTALLINE CELLULO SE (UNII: OP1R32D61U)		
POLYETHYLENE GLYCOL, UNSPECIFIED (UNII: 3WJQ0SDW1A)		
TALC (UNII: 7SEV7J4R1U)		
TITANIUM DIO XIDE (UNII: 15FIX9 V2JP)		

Product Characteristics				
Color	PINK (light pink)	Score	2 pieces	
Shape	OVAL	Size	16 mm	
Flavor		Imprint Code	CALAN;SR;180	
Contains				

ı	Packaging				
ı	#	Item Code	Package Description	Marketing Start Date	Marketing End Date
l	1 1	NDC:0025-1911-31	100 in 1 BOTTLE; Type 0: Not a Combination Product	12/15/1989	03/31/2021

Marketing Information			
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
NDA	NDA0 19 152	12/15/1989	03/31/2021

CALAN SR

verapamil hydrochloride tablet, film coated, extended release

Product Information			
Product Type	HUMAN PRESCRIPTION DRUG	Item Code (Source)	NDC:0025-1891
Route of Administration	ORAL		

Active Ingredient/Active Moiety			
Ingredient Name	Basis of Strength	Strength	
VERAPAMIL HYDRO CHLO RIDE (UNII: V38880EY5R) (VERAPAMIL - UNII:CJ0 O 37 KU 29)	VERAPAMIL HYDROCHLORIDE	240 mg	

Inactive Ingredients

Ingredient Name	Strength
CARNAUBA WAX (UNII: R12CBM0EIZ)	
HYPROMELLOSE, UNSPECIFIED (UNII: 3NXW29V3WO)	
MAGNESIUM STEARATE (UNII: 70097M6I30)	
MICRO CRYSTALLINE CELLULO SE (UNII: OP1R32D61U)	
POLYETHYLENE GLYCOL, UNSPECIFIED (UNII: 3WJQ0SDW1A)	
TALC (UNII: 7SEV7J4R1U)	
TITANIUM DIO XIDE (UNII: 15FIX9 V2JP)	
D&C YELLOW NO. 10 (UNII: 35SW5USQ3G)	
FD&C BLUE NO. 2 (UNII: L06K8R7DQK)	
ALUMINUM OXIDE (UNII: LMI26O6933)	

Product Characteristics				
Color	GREEN (light green)	Score	2 pieces	
Shape	OVAL (capsule shaped)	Size	19 mm	
Flavor		Imprint Code	CALAN;SR;240	
Contains				

ı	P	ackaging			
ı	#	Item Code	Package Description	Marketing Start Date	Marketing End Date
ı	1	NDC:0025-1891-31	100 in 1 BOTTLE; Type 0: Not a Combination Product	12/16/1986	

Marketing Information			
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
NDA	NDA0 19 152	12/16/1986	

CALAN SR

Product Information				
Product Type	HUMAN PRESCRIPTION DRUG	Item Code (Source)	NDC:0025-2712	
Route of Administration	ORAL			

Active Ingredient/Active Moiety			
Ingredient Name	Basis of Strength	Strength	
VERAPAMIL HYDRO CHLO RIDE (UNII: V38880EY5R) (VERAPAMIL - UNII:CJ0O37KU29)	VERAPAMIL HYDROCHLORIDE	120 mg	

Inactive Ingredients	
Ingredient Name	Strength
CARNAUBA WAX (UNII: R12CBM0EIZ)	

HYPROMELLOSE, UNSPECIFIED (UNII: 3NXW29V3WO)	
MAGNESIUM STEARATE (UNII: 70097M6I30)	
MICRO CRYSTALLINE CELLULO SE (UNII: OP1R32D61U)	
POLYETHYLENE GLYCOL, UNSPECIFIED (UNII: 3WJQ0SDW1A)	
TALC (UNII: 7SEV7J4R1U)	
TITANIUM DIO XIDE (UNII: 15FIX9 V2JP)	

Product Characteristics				
Color	PURPLE (light violet)	Score	no score	
Shape	OVAL	Size	11mm	
Flavor		Imprint Code	CALAN;SR;120	
Contains				

l	Packaging			
ı	# Item Code	Package Description	Marketing Start Date	Marketing End Date
ı	1 NDC:0025-2712-31	100 in 1 BOTTLE; Type 0: Not a Combination Product	09/17/2020	09/17/2020

Marketing Information			
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
NDA	NDA0 19 152	09/17/2020	09/17/2020

CALAN SR

Product Information				
Product Type	HUMAN PRESCRIPTION DRUG	Item Code (Source)	NDC:0025-2818	
Route of Administration	ORAL			

Active Ingredient/Active Moiety		
Ingredient Name	Basis of Strength	Strength
VERAPAMIL HYDRO CHLO RIDE (UNII: V38880EY5R) (VERAPAMIL - UNII:CJ0O37KU29)	VERAPAMIL HYDROCHLORIDE	180 mg

Inactive Ingredients		
Ingredient Name	Strength	
CARNAUBA WAX (UNII: R12CBM0 EIZ)		
HYPROMELLOSE, UNSPECIFIED (UNII: 3NXW29V3WO)		
MAGNESIUM STEARATE (UNII: 70097M6I30)		
MICRO CRYSTALLINE CELLULO SE (UNII: OP1R32D61U)		
POLYETHYLENE GLYCOL, UNSPECIFIED (UNII: 3WJQ0SDW1A)		
TALC (UNII: 7SEV7J4R1U)		

TITANIUM DIO XIDE (UNII: 15FIX9 V2JP)

Product Ch	Product Characteristics				
Color	PINK (light pink)	Score	2 pieces		
Shape	OVAL	Size	16 mm		
Flavor		Imprint Code	CALAN;SR;180		
Contains					

ı	Packaging			
	# Item Code	Package Description	Marketing Start Date	Marketing End Date
	1 NDC:0025-2818-31	100 in 1 BOTTLE; Type 0: Not a Combination Product	12/16/2020	

Marketing Information				
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date	
NDA	NDA0 19 152	12/16/2020		

CALAN SR

Product Information			
Product Type	HUMAN PRESCRIPTION DRUG	Item Code (Source)	NDC:0025-2924
Route of Administration	ORAL		

Active Ingredient/Active Moiety			
Ingredient Name	Basis of Strength	Strength	
VERAPAMIL HYDRO CHLO RIDE (UNII: V38880EY5R) (VERAPAMIL - UNII:CJ0 O 37 K U 29)	VERAPAMIL HYDROCHLORIDE	240 mg	

Inactive Ingredients			
Ingredient Name	Strength		
CARNAUBA WAX (UNII: R12CBM0EIZ)			
HYPROMELLOSE, UNSPECIFIED (UNII: 3NXW29V3WO)			
MAGNESIUM STEARATE (UNII: 70097M6I30)			
MICRO CRYSTALLINE CELLULO SE (UNII: OP1R32D61U)			
POLYETHYLENE GLYCOL, UNSPECIFIED (UNII: 3WJQ0SDW1A)			
TALC (UNII: 7SEV7J4R1U)			
TITANIUM DIO XIDE (UNII: 15FIX9 V2JP)			
D&C YELLOW NO. 10 (UNII: 35SW5USQ3G)			
FD&C BLUE NO. 2 (UNII: L06K8R7DQK)			
ALUMINUM OXIDE (UNII: LMI26O6933)			

Product Characteristics			
Color	GREEN (light green)	Score	2 pieces
Shape	OVAL (capsule shaped)	Size	19 mm
Flavor		Imprint Code	CALAN;SR;240
Contains			

ı	Packaging			
ı	# Item Code	Package Description	Marketing Start Date	Marketing End Date
ı	1 NDC:0025-2924-31	100 in 1 BOTTLE; Type 0: Not a Combination Product	09/17/2020	09/17/2020

Marketing Information			
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
NDA	NDA0 19 152	09/17/2020	09/17/2020

Labeler - Pfizer Laboratories Div Pfizer Inc (134489525)

Establishment			
Name	Address	ID/FEI	Business Operations
Pfizer Pharmaceuticals LLC		829084552	ANALYSIS(0025-1891, 0025-1901, 0025-1911, 0025-2712, 0025-2818, 0025-2924), MANUFACTURE(0025-1891, 0025-1901, 0025-1911, 0025-2712, 0025-2818, 0025-2924), PACK(0025-1891, 0025-1901, 0025-1911, 0025-2712, 0025-2818, 0025-2924), LABEL(0025-1891, 0025-1901, 0025-1911, 0025-2712, 0025-2818, 0025-2924)

Establishment			
Name	Address	ID/FEI	Business Operations
Pfizer Inc		943955690	ANALYSIS(0025-1891, 0025-1901, 0025-1911, 0025-2712, 0025-2818, 0025-2924)

Revised: 2/2021 Pfizer Laboratories Div Pfizer Inc